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(54) Title: DNA SEQUENCES, RECOMBINAN LINKED LYMPHOCYTE FUNCTION	T DN	A MOLECULES AND PROCESSES	FOR PRODUCING PI-
LINKED LYMPHOCYTE FUNCTION	A550	CIATED ANTIGEN-3	

(57) Abstract

Polypeptides that bind to CD2, the receptor on the surface of T-lymphocytes. Most preferably, the polypeptides are soluble. DNA sequences that code on expression and or secretion in appropriate unicellular hosts for those polypeptides. Methods of making and using those polypeptides in therapy and diagnosis.

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DNA SEQUENCES, RECOMBINANT DNA MOLECULES AND PROCESSES FOR PRODUCING PI-LINKED LYMPHOCYTE FUNCTION ASSOCIATED ANTIGEN-3

5 This invention relates to DNA sequences, recombinant DNA molecules and processes for producing Lymphocyte Function Associated Antigen-3 (PI-Linked form of LFA-3). More particularly, the invention relates to DNA sequences that are characterized in 10 that they code on expression in an appropriate unicellular host for a soluble PI-linked form of LFA-3 or derivatives thereof that bind to CD2, the receptor on the surface of T-lymphocytes. In accordance with this invention, unicellular hosts transformed with 15 these DNA sequences and recombinant DNA molecules containing them may also be employed to produce LFA-3 essentially free of other proteins of human origin. This novel antigen may then be used in the therapeutic and diagnostic compositions and methods of this 20 invention.

BACKGROUND OF THE INVENTION

T-lymphocytes play a major role in the immune response by interacting with target and antigen presenting cells. For example, the T-lymphocyte mediated killing of target cells is a multi-step process involving adhesion of a cytolytic T-lymphocyte to a target cell. And, helper T-lymphocytes initiate

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the immune response by adhesion to antigen-presenting cells.

These interactions of T-lymphocytes with target and antigen-presenting cells are highly specific and depend on the recognition of an antigen on the target or antigen-presenting cell by one of the many specific antigen receptors on the surface of the T-lymphocyte.

The receptor-antigen interaction of T-lymphocytes and other cells is also facilitated by 10 various T-lymphocyte surface proteins, e.g., the antigen receptor complex CD3(T3) and accessory molecules CD4, LFA-1, CD8, and CD2. It is also dependent on accessory molecules, such as LFA-3, ICAM-1 and MHC that are expressed on the surface of the target 15 or antigen-presenting cells. In fact, it is hypothesized that the accessory molecules on the T-lymphocytes and on the target or antigen-presenting cells interact with each other to mediate intercellular 20 adhesion. Accordingly, these accessory molecules are thought to enhance the efficiency of lymphocyteantigen-presenting cell and lymphocyte-target cell interactions and to be important in leukocyte-endothelial cell interactions and lymphocyte 25 recirculation.

For example, recent studies have suggested that there is a specific interaction between CD2 (a T-lymphocyte accessory molecule) and LFA-3 (a target cell accessory molecule) that mediates T-lymphocyte adhesion to the target cell. This adhesion is essential to the initiation of the T-lymphocyte functional response (M. L. Dustin et al., "Purified Lymphocyte Function-Associated Antigen-3 Binds To CD2 And Mediates T Lymphocyte Adhesion," J. Exp. Med., 165, pp. 677-92 (1987); Springer et al., "The Lymphocyte Function-Associated LFA-1, CD2, and LFA-3 Molecules: Cell Adhesion Receptors Of The Immune System", Ann.

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Rev. Immunol., 5, pp. 223-52 (1987)). And, monoclonal antibodies to either LFA-3 or CD2 have been shown to inhibit a spectrum of cytolytic T lymphocyte and helper T lymphocyte dependent responses (F. Sanchez-Madrid et al., "Three Distinct Antigens Associated With Human T-Lymphocyte-Mediated Cytolysis: LFA-1, LFA-2, And LFA-3", Proc. Natl. Acad. Sci. USA, 79, pp. 7489-93 (1982)).

LFA-3 is found on antigen-presenting cells, 10 and target cells, specifically on monocytes, granulocytes, CTL's, B-lymphoblastoid cells, smooth muscle cells, vascular endothelial cells, and fibroblasts (Springer et al., supra). LFA-3 exists as two distinct cell surface forms (Dustin et al., "Anchoring 15 Mechanisms For LFA-3 Cell Adhesion Glycoprotein At Membrane Surface", Nature, 329, pp. 846-848 (1987)). These forms differ mainly by their mechanism of attachment to lipid bilayers of biological membranes. One such anchoring mechanism is via a stretch of 20 hydrophobic amino acids, also referred to as a transmembrane domain, which penetrates the lipid bilayers. cDNA encoding this form of LFA-3, also referred to as an integrated membrane form, has been cloned and sequenced (B. Wallner et al., J.Exp.Med., 25-166, pp. 923-32 (1987)).

Alternatively, LFA-3 has been reported to insert into the membrane of B-lymphoblastoid cells via a phosphatidylinositol ("PI")-containing glycolipid covalently attached to the C-terminus of the protein (Dustin et al., supra). Membrane insertion of this type was deduced by observing the presence of protein after adding to the cell surface Phosphoinositol specific Phospholipase C. This enzyme releases only the PI-linked form of proteins. It does not affect the integrated membrane form. Thus, the release of LFA-3 in the presence of this enzyme suggests that LFA-3 has a PI-linked form.

The PI-linked form of LFA-3 is believed to be derived from alternative RNA splicing of a gene transcript. It appears to be selectively expressed in different cell types, and during different stages of development than the transmembrane form of LFA-3.

It would be desirable to obtain large amounts of a recombinant PI-linked form of LFA-3, than would be available from purification from natural sources, e.g. lymphoblastoid cells. More desirable would be to obtain large amounts of soluble LFA-3 from a PI-linked form of LFA-3.

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SUMMARY OF THE INVENTION

This invention solves these problems: aspect of this invention is the production of a recombinant PI-linked form of LFA-3. Another aspect 15 of this invention is the production of soluble LFA-3 from a PI-linked form of LFA-3. The latter embodiment is accomplished by expressing DNA sequences encoding a PI-linked form of LFA-3 in cell lines deficient in a PI-linkage attachment mechanism. A still further 20 aspect of this invention is the process of producing a soluble LFA-3 derived from a PI-linked form of LFA-3. This embodiment is accomplished by removing those portions of the DNA sequence encoding the hydrophobic transmembrane region of the PI-linked form of LFA-3. 25

This invention accomplishes each of these goals by providing DNA sequences coding on expression in an appropriate unicellular host for a PI-linked form of LFA-3 or derivatives thereof.

This invention also provides recombinant DNA molecules containing those DNA sequences and unicellular hosts transformed with them. Those hosts permit the production of large quantities of the PI-linked form of LFA-3, and its derivatives, of this invention for use in a wide variety of therapeutic and diagnostic compositions and methods.

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The DNA sequences of this invention are selected from the group consisting of:

- (a) the DNA sequence of the DNA insert carried in phage $\lambda P24$; and
- (b) DNA sequences which code on expression for a polypeptide coded for on expression by the foregoing DNA sequence.

The DNA sequences of this invention are also selected from derivatives of the DNA insert carried in phage $\lambda P24$ produced by removing those portions of the DNA sequence encoding the hydrophobic transmembrane region of the PI-linked form of LFA-3.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 depicts the amino acid sequences of the N-terminal and various peptide fragments of human LFA-3, purified from human erythrocytes using immunoaffinity chromatography.

Figure 2 depicts two pools of chemically synthesized oligonucleotide DNA probes derived from the amino acid sequence of a human LFA-3 purified from human erythrocytes.

Figure 3 depicts the DNA sequence of the DNA insert carried in phage $\lambda P24$ and the amino acid sequence deduced therefrom.

Figure 4 depicts the relevant portions of sequencing plasmid pNN01.

Figure 5 depicts the nucleotide sequence of probes LF-10, LF-11, NN-A, NN-B, NN-C, and NN-D.

Figure 6 depicts a comparison of the DNA insert carried in phage λP24 (and the deduced amino acid sequence) and the DNA insert carried in phage λHT16 (and the deduced amino acid sequence), which codes for an integrated membrane form of LFA-3.

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DETAILED DESCRIPTION OF THE INVENTION

We isolated the DNA sequences of this invention from a \(\lambda\)gt10 cDNA library derived from peripheral blood lymphocytes. However, we could also have employed libraries prepared from other cells that express a PI-linked form of LFA-3. These include, for example, monocytes, granulocytes, CTL's, B-lymphoblastoid cells, smooth muscle cells, endothelial cells and fibroblasts. We also could have used a human genomic bank.

For screening this library, we used a series of chemically synthesized anti-sense oligonucleotide DNA probes. We selected these probes from a consideration of the amino acid sequences of various fragments of LFA-3 that we determined using LFA-3 purified from human erythrocytes. These fragments are depicted in Figure 1. We selected amino acids from various areas that permitted the construction of oligonucleotide probes of minimal degeneracy.

We prepared two pools of probes: LFl and LF2-5. These pools are depicted in Figure 2. LFl is a 32-fold degenerate 20-mer and LF2-5 is a 384-fold degenerate 20-mer. Because of the high degeneracy of this latter pool, we subdivided the pool into four subpools -- LF2, LF3, LF4 and LF5 -- of 96-fold degeneracy each.

For screening, we hybridized our oligonucleotide probes to our cDNA libraries utilizing a
plaque hybridization screening assay. We selected
clone -- P24 -- hybridizing to one of our probes.
And, after isolating and subcloning the cDNA insert
of the selected clone, P24, into plasmids, we determined its nucleotide sequence and deduced the amino
acid sequence from those nucleotide sequences.

We have depicted in Figure 3 the nucleotide sequence of the cDNA insert of phage $\lambda P24$ and the amino acid sequence deduced therefrom. As shown in

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Figure 3, this cDNA insert has an open reading frame of 720 bp (240 amino acids), a 17 bp 5' untranslated region and a 93 bp 3' untranslated region. Also present is a transmembrane domain, i.e., from

N₆₆₂-N₇₂₅. The 3' untranslated region of P24 contains a poly(A) adenylation site. The P24 cDNA codes for a 240 amino acid protein, including a 28 amino acid signal sequence.

We have depicted in Figure 6 a comparison of the DNA sequences and deduced amino acid sequences of a membrane integrated form of LFA-3 (HT16) and the PI-linked form of LFA-3 of this invention. From this comparison, it is apparent that the last 14 amino acids (AA₂₀₉-AA₂₂₂) including a cytoplasmic domain at the C-terminus of the membrane integrated form of LFA-3 are replaced by 4 different amino acids in the PI-linked form of LFA-3.

The DNA sequences of this invention:

- (a) the DNA sequence of the DNA insert20 P24 carried in phage λP24; and
 - (b) DNA sequences which code on expression for a polypeptide coded for on expression by the foregoing DNA sequence,
- e.g., the cDNA sequence depicted in Figure 3 and contained in deposited clone λP24, may be used, as described below, in a variety of ways in accordance with this invention.

The DNA sequences, portions of them, or synthetic or semi-synthetic copies of them, may be used as a starting material to prepare various mutations. Such mutations may be either silent, i.e., the mutation does not change the amino acid sequence encoded by the mutated codon, or non-silent, i.e., the mutation changes the amino acid sequence encoded by the mutated codon. Both types of mutations may be advantageous in producing or using the LFA-3's of this invention. For example, these

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mutations may permit higher levels of production, easier purification, or production of secreted shortened or soluble forms of PI-linked forms of LFA-3.

5 The DNA sequences of this invention are also useful for producing the PI-linked forms of LFA-3, or its derivatives, coded on expression by them in unicellular hosts capable of attaching proteins by PI-linkage, e.g., CHO cells, transformed with those DNA sequences. Preferably, according to 10 a second embodiment of this invention, these DNA sequences may be expressed in a cell line deficient in the PI-linkage attachment mechanism, such as mouse L-cells, e.g., L-M (tk) cells. In this case the LFA-3 may be secreted into the medium in a soluble 15 This secreted form of LFA-3 is approximately 3 kd smaller than other forms of LFA-3 retained intracellularly in L-M (tk) cells or extracted from CHO cells. While not wishing to be bound by theory, 20 we believe that the DNA sequences of the present invention produce and secrete a smaller soluble form of LFA-3 because a portion of the transmembrane region is cleaved before or after secretion by cells deficient in a PI-linkage attachment mechanism, and there-25 fore efficient attachment of the PI-linked form of LFA-3 to the cell surface is prevented.

According to another embodiment of this invention DNA sequences encoding a PI-linked form of LFA-3 may be modified as compared to that of Figure 3 (amino acids -28 to 212) to remove from it portions that code for the hydrophobic transmembrane region, e.g., from about nucleotide 662 to 731, to allow production of soluble LFA-3 protein in any cell transformed with those modified sequences.

As well known in the art, the DNA sequences of this invention are expressed by operatively linking them to an expression control sequence in an appro-

priate expression vector and employed in that expression vector to transform an appropriate unicellular host.

Such operative linking of a DNA sequence of this invention to an expression control sequence, 5 of course, includes the provision of a translation start signal in the correct reading frame upstream of the DNA sequence. If the particular DNA sequence of this invention being expressed does not begin with a methionine, e.g., a mature PI-linked form of 10 LFA-3 which begins with a phenylalanine, the start signal will result in an additional amino acid -methionine -- being located at the N-terminus of the product. While such methionyl-containing-product may be employed directly in the compositions and 15 methods of this invention, it is usually more desirable to remove the methionine before use. Methods are available in the art to remove such N-terminal methionines from polypeptides expressed with them. For example, certain hosts and fermenta-20 tion conditions permit removal of substantially all of the N-terminal methionine in vivo. Other hosts require in vitro removal of the N-terminal methionine. However, such $\underline{\text{in}}$ $\underline{\text{vivo}}$ and $\underline{\text{in}}$ $\underline{\text{vitro}}$ methods are well known in the art. Furthermore, the LFA-3's of this 25 invention may include amino acids in addition to the N-terminal methionine at the N-terminus. may be used with those amino acids or they may be cleaved with the N-terminal methionine before use. 30

A wide variety of host/expression vector combinations may be employed in expressing the DNA sequences of this invention. Useful expression vectors, for example, may consist of segments of chromosomal, non-chromosomal and synthetic DNA sequences, such as various known derivatives of SV40 and known bacterial plasmids, e.g., plasmids from E.coli including col E1, pCR1, pBR322, pMB9 and their

derivatives, wider host range plasmids, e.g., RP4, phage DNAs, e.g., the numerous derivatives of phage λ, e.g., NM989, and other DNA phages, e.g., M13 and Filamenteous single stranded DNA phages, yeast plasmids such as the 2μ plasmid or derivatives thereof, vectors useful in eukaryotic cells, such as vectors useful in animal cells and vectors derived from combinations of plasmids and phage DNAs, such as plasmids which have been modified to employ phage DNA or other expression control sequences. In the preferred embodiments of this invention, we employ BG8, a pBR312-related vector [R. Cate et al., Cell, 45, pp. 685-98 (1986)].

In addition, any of a wide variety of expression control sequences -- sequences that con-15 trol the expression of a DNA sequence when operatively linked to it -- are used in these vectors to express the DNA sequence of this invention. Such useful expression control sequences, include, for example, the early and late promoters of SV40 or the 20 adenovirus, the <u>lac</u> system, the <u>trp</u> system, the <u>TAC</u> or TRC system, the major operator and promoter regions of phage λ , the control regions of fd coat protein, the promoter for 3-phosphoglycerate kinase or other glycolytic enzymes, the promoters of acid phosphatase, 25 e.g., Pho5, the promoters of the yeast α -mating factors, and other sequences known to control the expression of genes of prokaryotic or eukaryotic cells or their viruses, and various combinations thereof. For animal cell expression (e.g., L-M (tk⁻) 30 cells), we prefer to use an expression control sequence derived from the major late promoter of adenovirus 2.

A wide variety of unicellular host cells

are also useful in expressing the DNA sequences of
this invention. These hosts may include well known
eukaryotic and prokaryotic hosts, such as strains of

E.coli, Pseudomonas, Bacillus, Streptomyces, fungi, such as yeasts, and animal cells, such as CHO and RI.1, B-W and L-M cells, African green monkey cells, such as COS1, COS7, BSC1, BSC40, and BMT10, and human cells and plant cells in tissue culture. For expression of a soluble form of LFA-3, an appropriate host cell is defective in the PI attachment of proteins. We prefer L-M (tk⁻) cells.

It should of course be understood that not all vectors, expression control sequences and hosts 10 will function equally well to express the DNA sequences of this invention. Neither will all hosts function equally well with the same expression system. However, one of skill in the art may make a selection 15 among these vectors, expression control sequences, and hosts without undue experimentation and without departing from the scope of this invention. For example, in selecting a vector, the host must be considered because the vector must replicate in it. The vector's copy number, the ability to control 20 that copy number, and the expression of any other proteins encoded by the vector, such as antibiotic markers, should also be considered.

In selecting an expression control sequence, 2.5 a variety of factors should also be considered. These include, for example, the relative strength of the system, its controllability, and its compatibility with the particular DNA sequence, of this invention, particularly as regards potential secondary structures. Unicellular hosts should be selected by 30 consideration of their compatibility with the chosen vector, the toxicity of the product coded on expression by the DNA sequences of this invention to them, their secretion characteristics, their ability to 35 fold proteins correctly, their fermentation requirements, and the ease of purification of the products

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coded on expression by the DNA sequences of this invention.

Within these parameters one of skill in the art may select various vector/expression control system/host combinations that will express the DNA sequences of this invention on fermentation or in large scale animal culture, e.g., mouse cells or CHO cells.

The polypeptides produced on expression of
the DNA sequences of this invention may be isolated
from the fermentation or animal cell cultures and
purified in a variety of ways well known in the art.
Such isolation and purification techniques depend on
a variety of factors, such as how the product is
produced, whether or not it is soluble or insoluble,
and whether or not it is secreted from the cell or
must be isolated by breaking the cell. One of skill
in the art, however, may select the most appropriate
isolation and purification techniques without
departing from the scope of this invention.

The polypeptides of this invention are useful in compositions and methods to block or to augment the immune responses. For example, they are active in inhibiting cytolytic T-lymphocyte activity by interfering with T-cell interaction with target cells. They have a similar blocking or augmenting effect on the immune response because they interfere with the interaction of helper T-cells and target cells. Furthermore, the compounds of this invention may be used to target specific T cells for lysis and immune suppression or to deliver drugs, such as lymphokines, to the specifically targeted T-cells. More preferably, soluble derivatives of the polypeptides of this invention may be employed to saturate the CD2 sites of T-lymphocytes thus inhibiting T-cell activation. This is plainly of great utility in graft-vs-host disease, in autoimmune diseases, e.g.,

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rheumatoid arthritis, and in preventing allograft rejection. Furthermore, the polypeptides of this invention are preferred over monoclonal antibodies to a PI-linked form of LFA-3 or CD2 because the polypeptides of this invention are less likely to elicit immune responses in humans than are antibodies raised in species other than humans. The therapeutic compositions of this invention typically comprise an immunosuppressant or enhancement effective amount of such polypeptide and a pharmaceutically acceptable carrier. The therapeutic methods of this invention comprise the steps of treating patients in a pharmaceutically acceptable manner with those compositions.

in these therapies may be in a variety of forms.

These include, for example, solid, semi-solid and liquid dosage forms, such as tablets, pills, powders, liquid solutions or suspensions, liposomes, suppositories, injectable and infusable solutions. The preferred form depends on the intended mode of administration and therapeutic application. The compositions also preferably include conventional pharmaceutically acceptable carriers and adjuvants which are known to those of skill in the art.

Preferably, the compositions of the invention are in the form of a unit dose and will usually be administered to the patient one or more times a day.

Generally, the pharmaceutical compositions of the present invention may be formulated and administered using methods and compositions similar to those used for other pharmaceutically important polypeptides (e.g., alpha-interferon). Thus, the polypeptides may be stored in lyophilized form, reconstituted with sterile water just prior to administration, and administered by the usual routes of administration such as parenteral, subcutaneous, intravenous or intralesional routes.

The polypeptides of this invention or antibodies against them are also useful in diagnostic
compositions and methods to detect T-cell subsets or
CD2+ cells or to monitor the course of diseases
characterized by excess or depleted T-cells, such as
autoimmune diseases, graft versus host diseases and
allograft rejection. Still further, the polypeptides
of this invention may be used to screen for inhibitors
of LFA-3 mediated adhesion useful for inhibiting
activation of T lymphocytes or T lymphocyte mediated
killing of target cells. Such screening techniques
are well-known in the art.

Finally, the polypeptides of this invention or antibodies against them are useful in separating B and T cells. For example, when bound to a solid support the polypeptides of this invention or antibodies to them will separate B and T cells.

In order that this invention may be better understood, the following examples are set forth.

These examples are for purposes of illustration only, and are not to be construed as limiting the scope of the invention in any manner.

Synthesis Of Oligonucleotide Probes

We obtained a sample of LFA-3 (Dana Farber Cancer Institute, Boston, Massachusetts) previously 25 purified as described by M. Dustin et al., J. Exp. Med., supra and sequenced as described by B. Wallner et al., supra. Next, we chemically synthesized two pools of anti-sense oligonucleotide DNA probes coding for regions from the amino terminal sequence of our 30 sample of LFA-3 characterized by minimal nucleic acid degeneracy (see underscoring in Figure 1) on an Applied Biosystems 30A DNA synthesizer. For each selected amino acid sequence, we synthesized pools of probes complementary to all possible codons. We 35 synthesized the probes anti-sense to enable hybrid-

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ization of them to the corresponding sequences in DNA as well as in mRNA. We labelled our oligonucleotide probes using $[\gamma^{-32}P]$ -ATP and polynucleotide kinase (Maxam and Gilbert, <u>Proc. Natl. Acad. Sci.</u>, 74, p. 560 (1977)).

As depicted in Figure 2, the oligonucleotide probe pool LF1 was a 20-mer with 32-fold degeneracy. Probe pool LF2-5, was a 20-mer with 384-fold degeneracy. However, to reduce its degeneracy, we synthesized this pool in four subpools of 96-fold degeneracy each by splitting the degenerate codon for Gly into one of its four possible nucleotides for each subpool. We then selected the subpool containing the correct sequence from the three pools containing incorrect sequences by hybridization of the individual subpools to Northern blots containing human tonsil mRNA, as described previously (Wallner et al., Nature, 320, pp. 77-81 (1986)). Oligonucleotide probe subpool LF2 hybridized to a 1300 nucleotide transcript in human tonsil RNA, which suggested that it contained the correct sequence. Hence, we used it and pool LF1 for screening our various libraries.

Construction Of Agt10 Peripheral Blood Lymphocytes cDNA Library

25 To prepare our Peripheral Blood Lymphocytes (PBL) DNA library, we processed PBL from leukophoresis #9 through one round of absorption to remove monocytes. We then stimulated the non-adherent cells with IFN- γ 1000 U/ml and 10 μ g/ml PHA for 24 h. We isolated RNA from these cells using phenol extraction 30 (Maniatis et al., Molecular Cloning, p. 187 (Cold Spring Harbor Laboratory) (1982)) and prepared poly A mRNA by one round of oligo dT cellulose chromatography. We ethanol precipitated the RNA, dried it in a speed vac and resuspended the RNA in 10 μ 1 $\rm H_2O$ 35 (0.5 $\mu g/\mu l$). We treated the RNA for 10 min at room temperature in CH_3HgOH (5mM final concentration) and

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β-mercaptoethanol (0.26 M). We then added the methyl mercury treated RNA to 0.1 M Tris-HCl (pH 8.3) at 43°C, 0.01 M Mg, 0.01 M DTT, 2 mM Vanadyl complex, 5 μg oligo dT_{12-18} , 20 mM KCl, 1 mM dCTP, dGTP, dTTP, 0.5 mM dATP, 2 μCi[α - 32 P]dATP and 30 U 1.5μl AMV reverse transcriptase (Seikagaku America) in a total volume of 50 μl. We incubated the mixture for 3 min at room temperature and 3 h at 44°C after which time we stopped the reaction by the addition of 2.5 μl of 0.5 M EDTA.

We extracted the reaction mixture with an equal volume of phenol:chloroform (1:1) and precipitated the aqueous layer two times with 0.2 volume of 10 M $\rm NH_4OAc$ and 2.5 volumes EtOH and dried it under vacuum. The yield of cDNA was 1.5 μg .

We synthesized the second strand according to the methods of Okayama and Berg (Mol. Cell. Biol., 2, p. 161 (1982)) and Gubler and Hoffman (Gene, 25, p. 263 (1983)), except that we used the DNA polymerase I large fragment in the synthesis.

We blunt ended the double-stranded cDNA by resuspending the DNA in 80 μ l TA buffer (0.033 M Tris Acetate (pH 7.8); 0.066 M KAcetate; 0.01M MgAcetate; 0.001M DTT; 50 μ g/ml BSA), 5 μ g RNase A, 4 units RNase H, 50 μ M β NAD , 8 units E.coli ligase, 0.3125 mM dATP, dCTP, dGTP, and dTTP, 12 units T₄ polymerase and incubated the reaction mixture for 90 min at 37°C, added 1/20 volume of 0.5M EDTA, and extracted with phenol:chloroform. We chromatographed the aqueous layer on a G150 Sephadex column in 0.01M Tris-HCl (pH 7.5), 0.1 M NaCl, 0.001 M EDTA and collected the lead peak containing the double-stranded cDNA and ethanol precipitated it. Yield: 605 μ g cDNA.

We ligated the double-stranded cDNA to

35 linker 35/36

5'AATTCGAGCTCGAGCGCGGCCGC3'

3' GCTCGAGCTCGCGCCGGCG5'

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using standard procedures. We then size selected the cDNA for 800 bp and longer fragments on a S500 Sephacryl column, and ligated it to EcoRI digested Agt10. We packaged aliquots of the ligation reaction in Gigapak (Strategene) according to the manufacturer's protocol. We used the packaged phage to infect E.coli BNN102 cells and plated the cells for amplification. The resulting library contained 1.125x10⁶ independent recombinants.

10 Screening Of The Libraries

We screened the PBL cDNA library prepared above with our labelled oligonucleotide probe LF1 using the plaque hybridization screening technique of Benton and Davis (Science, 196, p. 180 (1977)).

We pelleted an overnight culture of BNN102 cells in L broth and 0.2% maltose and resuspended it in an equal volume of SM buffer (50 mM Tris-HCl (pH 7.5), 100 mM NaCl, 10 mM MgSO₄, and 0.01% gelatin). Thereafter, we preabsorbed 9 ml of cells with 1.5x10⁶ phage particles at room temperature for 15 minutes and plated them on 30 LB Mg plates.

After incubation at 37°C for 8 hours, we absorbed phages onto filters from the plates and lysed the filters by placing them onto a pool of 0.5 N NaOH/1.5 M NaCl for 5 minutes, and then submerged them for 5 min in the same buffer. We neutralized the filters by submerging them in 0.5 M Tris-HCl (pH 7.4), 1.5 M NaCl, two times for 5 minutes each, and rinsed them for 2 minutes in 1 M NH₄OAc, air dried the filters, and baked them for 2 hours at 80°C.

We prehybridized and hybridized the filters to oligonucleotide probe LF1 in 0.2% polyvinylpyrolidone, 0.2% ficoll (MW 400,000), 0.2% bovine serum albumin, 0.05 M Tris-HCl (pH 7.5), 1 M sodium chloride, 0.1% sodium pyrophosphate, 1% SDS, and 10%

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dextran sulfate (MW 500,000). We detected the hybridizing λ -cDNA sequences by autoradiography.

We initially selected 26 positive phages from the PBL library and rescreened these clones and plaque purified them at lower density using the same probe.

Sequencing Of The P24 cDNA Clone

We characterized the cDNA from a clone, P24, screened above by DNA sequencing analysis. We subcloned the NotI digested DNA from clone λP24 into vector pNN01 to give p24 and to facilitate sequence analysis.* The entire insert of λP24 is contained on a single NotI fragment. For subcloning, we used the vector's EcoRI site or SmaI site employing techniques in common use.

We determined the DNA sequences of our subclones largely by the method of Maxam and Gilbert (Meth. Enzymology, 65, pp. 499-560, (1980)). However, for some fragments, we used the related procedure of Church and Gilbert (Proc. Natl. Acad. Sci. USA, 81, p. 1991 (1984)). The structure of pNNO1 enables sequencing, by the Church-Gilbert approach, of the ends of an inserted fragment using NotI digestion and four 20-nucleotide long probes: NN-A, NN-B, NN-C and NN-D. See Figure 5.

Figure 3 shows the DNA sequence of the cDNA insert of phage $\lambda P24$. It also depicts the amino acid sequence deduced therefrom.

^{*} We constructed sequencing plasmid pNN01 by removing the synthetic polylinker of pUC8 by restriction digestion and replacing it with a new synthetic segment. The 2.5 kb backbone common to the pUC plasmids, which provides an origin of replication and confers ampicillin resistance, is unchanged. The novel synthetic portion of pNN01 is shown in Figure 4.

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Determination Of Linkage Form Of LFA-3 From P24 cDNA

We decided to characterize the linkage form of LFA-3 coded for by P24 cDNA. We choose the R1.1 cell line because it is known to express surface antigens that are attached to the membrane by a PI linkage.

We incubated 5 x 10⁶ cells of clones P24/R1.1, HT16/R1.1 (an R1.1 cell line transfected with cDNA coding for a membrane integrated form of LFA-3 (see, B. Wallner et al., <u>supra</u>)) and R1.1 cells with .5 µl of Phosphoinositol specific Phospholipase C (PIPLC) at 37°C for 1 hour. It is known that PIPLC upon incubation releases PI-linked proteins from the cell surface, while it has no effect on proteins attached to the cell surface by other mechanisms such as membrane integrated proteins (M. Low, <u>J. Biochem.</u>, 244, p. 1 (1987)).

from the cell surface of P24/R1.1 or HT16/R1.1 by the decrease of surface fluorescence assayed on FACS. We found that incubation of P24/R1.1 cells with PIPLC resulted in the release of 95% of surface LFA-3 while PIPLC did not have any effect on the fluorescence of R1.1 cells or HT16/R1.1 cells. This indicates that P24 cDNA codes for the PI-linked form of LFA-3.

Adhesion Of a PI-linked form of LFA-3 From P24/R1.1 To Other Cells

We next tested whether a PI-linked form of LFA-3 from P24 cDNA as expressed in R1.1 cells would mediate adherence of P24/R1.1 to other cells. We tested this by rosetting analysis with L-cells expressing CD2 cDNA (L114). We grew control L cells and L114 (CD2 transfected) cells, in a 9.6 cm² well of a 6 well tissue culture plate at a cell density

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of 3 \times 10⁵ cells per well. After washing the wells twice with Roswell Park Memorial Institute (RPMI 1060) medium to remove cell debris and dead cells, 1.5 \times 10^7 P24/R1.1 or R1.1 cells as a control were added per well. Plates were spun at 400 rpm for 2 minutes in a Sorvall Centrifuge at 4°C. After the cells were incubated at 4°C for 2 hours, the wells were washed with RPMI 1060 medium to remove excess P24/R1.1 or R1.1 cells. P24/R1.1 cells rosetted with the L114 cells as determined under the microscope. We 10 observed rosetting of P24/R1.1 with L114 cells but not with the untransfected control cells. This rosetting could be inhibited with MAb to LFA-3 (TS2/9) or MAb to CD2 (TS2/18). This indicates that a PIlinked form of LFA-3 is expressed on cell surface of 15 R1.1 cells in a conformation that allows interaction with recombinant CD2 expressed on mouse L-cells. P24/R1.1 cells or untransfected R1.1 cells do not rosette with untransfected mouse L-cells, indicating

Expression Of PI-linked Form Of LFA-3 From P24 cDNA In CHO cells

We inserted a Klenow blunt-ended NotI PIlinked form of LFA-3 cDNA fragment of p24 into a blunt-ended SalI site of plasmid pJOD-s to give pJOD-s-LFA3P24.

the specificity of these cellular interactions.

Vector pJOD-s has been deposited in the In Vitro International, Inc. Culture Collection, 611 P. Hammonds Ferry Rd., Linthicum, Maryland, 21090 on July 22, 1988 and has been assigned accession number 10179.

We next linearized pJOD-s-LFA3P24 with PvuI for transfection of CHO cells. We incubated 10µg of PvuI linearized DNA with 0.125 M CaCl₂ in TE and 1 x HEBS (137 mM NaCl, 5mM KCl, 0.0030 M Na₂HPO₄, .7 H₂O, 6mM Dextrose, 20 mM Hepes (pH 7.1)) at room

temperature for 20 minutes. DNA was added to cells in alpha⁺-MEM medium and incubated at 37°C for 4 hours. After removing the medium, cells were incubated at room temperature for 4 minutes in alpha⁺-MEM + 10% glycerol. Cells were rinsed with medium and grown for 2 days in alpha⁺-MEM, then transferred to selective medium (alpha⁻-MEM).

transferred to selective medium (alpha -MEM). We determined expression of a PI-linked form of LFA-3 by FACS analysis. To analyze by FACS, 1 x 10⁶ cells per each P24-CHO methotrexate clone and 10 control CHO cells were removed from the tissue culture dishes by incubation with Hank's BSS buffer, .5 M EDTA at 4°C for 15 minutes. The detached cells were then pelleted, resuspended in 50 μl of PBN buffer (1 x PBS, .5% BSA, .1% sodium azide) and incu-15 bated with 100 μl of MAb TS2/9 (1.2 mg/ml) (a gift of Tim Springer) on ice for 45 minutes. We next washed the cells two times with 1 ml PBN buffer and pelleted by centrifugation. The cell pellets were resuspended in 100µl of a 1:50 dilution of FCI 20 (Fluorescein Conjugated Affinity Purified F (ab') 2 Fragment Sheep Anti-Mouse IgG (Cappel, Biomedical, Pennsylvania)) in PBN buffer and incubated on ice for 30 minutes. Cells were pelleted by centrifugation and excess FCI was removed by resuspending the 25 cell pellets twice in 1 ml PBN buffer. We then resuspended the cells in 800 μl of 1 x PBS and determined the fluorescence intensity on FACS. We observed five clones showed between 5 to 50 fold higher 30 fluorescence than control CHO cells.

Expression Of A PI-linked Form Of LFA-3 From P24 cDNA In R1.1 cells

We used expression vector BG24 derived from expression vector BG312. BG24 was constructed by digesting plasmid p24 DNA with NotI and blunt-ended with Klenow. We next isolated a 860 bp NotI fragment

of p24 followed by ligation with an EcoRI linearized, blunt-ended expression vector BG368. BG368 was constructed as follows. Animal expression vector pBG312 (R. Cate et al., Cell, 45, pp. 685-98 (1986)) was

digested with EcoRI and BglII to delete one of each of the two EcoRI and the two BglII restriction sites (the EcoRI site at position 0 and the BglII site located at approximately position 900).

90 μg DNA of BG24 was linearized with NruI,
10 and cotransfected with 10 μg of NruI linearized pTCF
DNA (F. Grosveld et al., Nucleic Acid Res., 10,
p. 6715 (1982)) and 300 μg sonicated salmon sperm
DNA by DNA electroporation using a BIORAD (Richmond,
California) gene pulse at 0.29 UV with capacitance
15 set at 960 μFD. We selected for transfection R1.1
cells in RPMI 1060 medium + 1 mg/ml G418. We selected
single clones after limiting dilutions to 10³ cells
per well in a 96 well dish in selective medium.
Eight clones, resistant to G418, were assayed for a
PI-linked form of LFA-3 expression by FACS analysis as
described above.

described above. All eight P24/R1.1 clones expressed PI-linked form of LFA-3 at a level 10 to 1000 fold above R1.1 control cells.

Expression Of A PI-linked Form Of LFA-3 P24 cDNA In L Cells

To express our P24 cDNA in mouse L cells, we cotransfected 90 µg of plasmid BG24 DNA ,as described above, that was linearized with Nrul with 10µg plasmid pOPF DNA carrying a thymidine kinase gene (tk) (Grosveld et al., supra), linearized with Scal into 1 x 10⁷ L-M (tk) cells (C. P. Terhorst, J. Immun., 131, p. 2032 (1983)) by electroporation as described above. We selected for transfected cells by tk expression by growing them in DMEM + HAT at cell densities of 1 x 10⁵ cells per 100 mm plate. Clones were picked and expanded to 5 x 10⁵ cells per

100 mm dish to assay for expression of a PI-linked form of LFA-3 by FACS analysis as desribed above. We observed some expression at levels above control cells, although 70% of the PI-linked form of LFA-3 was secreted into the medium as discussed below.

Secretion Of LFA-3 From P24/L Cells

We further wanted to test whether P24/L cells secrete LFA-3 because this mouse L cell line --L-M(tk) -- is known to be deficient in a PI linkage 10 attachment mechanism. P24/L cells were metabolically labeled with 35 s-met and the 35 s-labelled PI-linked form of LFA-3 was precipitated from the medium with MAb TS2/9 (a gift of Tim Springer) as follows. 3 x10⁵ P24/L, HT16/L (B. Wallner et al., supra) or L(tk⁻) 15 cells were plated in 1 well each of a 6 well cell culture plate, grown overnight in DMEM-HAT complete medium (DMEM + HAT + 10% FCS + glutamine). Wells were than rinsed with 1x Minimal Essential Medium Eagle (modified) methionine free (MEM). For 35 S met 20 labeling, we added 1.5 ml of MEM medium (methionine free), plus glutamine, 2.5% complete DMEM and 225 $\mu\text{Ci}^{35}\text{S}$ met (New England Nuclear, Delaware, 1135 mCi/μm) to each well and incubated at 37°C for-18 hours. To 0.7 ml of medium 10 μ l of MAb TS2/9 25 coupled to agarose was added, and the mixture rocked at $4\,^{\circ}\text{C}$ overnight. To each well we added 300 μl of DOC buffer (20 mM Tris (pH 7.3), 50mM sodium chloride, .5% dioxycholate, .5% Triton x 100), scraped the cells off the plates, transferred to Eppendorf tubes, 30 vortexed and centrifuged for 15 minutes at room temperature. To 100 μl of the supernatant, 10 μl of MAb TS2/9 coupled to agarose was added and incubated overnight at 4°C with rocking. The TS2/9-agarose 35S-LFA-3 complex was pelleted by centrifuation, 35 washed three times with 1 ml of DOC buffer, and

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resuspended in 50 µl SDS-loading buffer. ³⁵S-LFA-3 (PI-linked form) was dissociated from TS2/9-agarose by heating the complex to 65°C for 10 minutes. The TS2/9 agarose was precipitated by centrifugation and 25 µl of the supernatant was electrophoresed on a reducing SDS-polyacrylamide gel. We observed precipitation of the 55 kd ³⁵S-labelled protein with MAb TS2/9 only from the medium of P24/L cells and not from medium of L(tk-) control cells.

We determined by SDS-PAGE that the ³⁵S-labeled LFA-3, secreted from P24/L cells is approximately 3 kd smaller than the ³⁵S-labelled LFA-3 retained intracellularly in P24/L cells or HT16/L cells. This indicates that a portion or all of the hydrophobic potential transmembrane region is removed before secretion, which prevents the efficient integration of a PI-linked form of LFA-3 into the cell surface membrane.

We deposited the following plasmid carrying
a PI-linked form of LFA-3 cDNA sequence of this invention in the In Vitro International, Inc. Culture
Collection in Linthicum, Maryland, on July 22, 1988:
p24

The plasmid has been assigned accession number IVI-10180.

While we have herein before presented a number of embodiments of this invention, it is apparent that our basic construction can be altered to provide other embodiments which utilize the processes and compositions of this invention. Therefore, it will be appreciated that the scope of this invention is to be defined by the claims appended hereto rather than the specific embodiments which we have presented by way of example.

CLAIMS:

- 1. A DNA sequence selected from the group consisting of:
- (a) the DNA sequence of the DNA insert carried in phage $\lambda P24$; and
- (b) DNA sequences which code on expression for a polypeptide coded for on expression by the foregoing DNA sequence.
- 2. A DNA sequence selected from the group consisting of a DNA sequence of the formula N_{1-830} of Figure 3, a DNA sequence of the formula N_{18-830} of Figure 3, a DNA sequence of the formula $N_{102-830}$ of Figure 3 and DNA sequences that code on expression for a polypepdite coded for on expression by any of the above DNA sequences.
- consisting of a DNA sequence of the formula N_{1-653}^{-1} $N_{738-830}^{-1}$ of Figure 3, a DNA sequence of the formula $N_{102-653}^{-1}$ $N_{738-830}^{-1}$ of Figure 3, a DNA sequence of the formula N_{1-662}^{-1} $N_{738-830}^{-1}$ of Figure 3, a DNA sequence of the formula N_{1-662}^{-1} $N_{738-830}^{-1}$ of Figure 3, a DNA sequence of the formula $N_{102-662}^{-1}$ $N_{738-830}^{-1}$ of Figure 3, a DNA sequence of the formula N_{1-638}^{-1} $N_{102-638}^{-1}$ $N_{102-638}^{-1}$ of Figure 3, a DNA sequence of the formula N_{1-701}^{-1} $N_{102-701}^{-1}$ $N_{102-701}^{-1}$ $N_{102-701}^{-1}$ $N_{102-701}^{-1}$ of Figure 3, a DNA sequence of the formula $N_{102-701}^{-1}$ $N_{102-701}^{-1}$
- 4. A recombinant DNA molecule comprising a DNA sequence according to claim 1 or 2, said DNA sequence being operatively linked to an expression control sequence in said recombinant DNA molecule.

- 5. A recombinant DNA molecule comprising a DNA sequence of claim 3, said DNA sequence being operatively linked to an expression control sequence in said recombinant DNA molecule.
- 6. The recombinant DNA molecule according to claim 4 or 5, wherein said expression control sequence is selected from the group consisting of the early or late promoters of SV40 or adenovirus, the <u>lac</u> system, the <u>trp</u> system, the <u>TAC</u> system, the <u>TRC</u> system, the major operator and promoter regions of phage λ , the control regions of fd coat protein, the promoter for 3-phosphoglycerate kinase or other glycolytic enzymes, the promoters of acid phosphatase and the promoters of the yeast α -mating factors.
- 7. The recombinant DNA molecule according to claim 4, wherein the molecule is selected from the group consisting of pJOD-s-LFA3P24 and BG24.
- 8. A unicellular host transformed with a recombinant DNA molecule according to claim 4.
- 9. A unicellular host transformed with a recombinant DNA molecule according to claim 4, characterized in that it is deficient in a PI-linkage attachment mechanism.
- 10. A unicellular host transformed with a recombinant DNA molecule according to claim 5.
- 11. The host according to claim 8 or 10 wherein said host is selected from the group consisting of strains of <u>E.coli</u>, <u>Pseudomonas</u>, <u>Bacillus</u>, <u>Streptomyces</u>, yeast, fungi, animal cells, plant cells and human cells in tissue culture.

- 12. The unicellular host according to claim 11, wherein the animal cell is selected from the group consisting of CHO and R1.1.
- 13. The unicellular host according to claim 9, wherein the host is L-M(tk⁻).
- 14. A method of producing a polypeptide comprising the steps of culturing a unicellular host according to claim 8.
- 15. The method according to claim 14 wherein the transformed host is selected from the group consisting of CHO(pJOD-s-LFA3P24) and R1.1(BG24).
- 16. A method of producing a soluble polypeptide comprising the step of culturing a unicellular host according to claim 9.
- 17. A method of producing a soluble polypeptide comprising the step of culturing a unicellular host according to claim 10.
- 18. The method according to claim 16 wherein the transformed host is selected from the group consisting of L-M(tk⁻)(BG24) and L-M(tk⁻)(pJOD-s-LFA3P24).
- 19. A polypeptide coded on expression by a DNA sequence selected from the group consisting of the DNA sequences of claim 1 or 2, said polypeptide being essentially free of other proteins of human origin.
- 20. A pharmaceutical composition comprising an immunosuppressant or enhancement effective

amount of a polypeptide of claim 19 and a pharmaceutically acceptable carrier.

- 21. A method of treating patients comprising the step of treating them in a pharmaceutically acceptable manner with a composition of claim 20.
- T-cell subsets, CD2+ cells or to monitor the course of diseases characterized by excess or depleted T-cells comprising a diagnostic effective amount of a polypeptide of claim 19 or an antibody thereto.
- 23. A method of detecting T-cell-subsets, CD2+ cells or for monitoring the course of diseases characterized by excess or depleted T-cells comprising the step of employing as a diagnostic a composition of claim 22.
- 24. A means for detecting T-cell subsets, CD2+ cells or for monitoring the course of diseases characterized by excess or depleted T-cells comprising a composition of claim 23.
- 25. A method of isolating an LFA-3 inhibitor said method comprising:
 - (a) transfecting a cell line with p24,
 - (b) conducting a binding assay with Jurkatt cells and said transfected cell line, and
 - (c) selecting molecules that inhibit binding of Jurkatt cells to the cells of step (a).

FIG. I

NH₂ Terminus : FSOOIYGVVYGXVTFHVPSNVP LFA-3 LKEVLWKKOKDKVAEL

T₇₂₋₇₃ : DKVAELENSEF

T₉; : VYLDTVSGSLTIYNLTS

T₁₀₅ : FFLYVLESLPSPTLTCAL

T₆₈ : GLIMYS

FIG. 5

LF-10 : cgtcgctcccagcaaccatggctcgtcggg
LF-11 : catggaaagttacattcccatacacaccc
NN-A : gatcctcacatcccaatccg
NN-B : tccaaccaccaatctcaaag
NN-C : cggattgggatgtgaggatc
NN-D : ctttgagattggtggttgga

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FIG. 2

Oligonucleotide Probe Pool LF1:

20mer, 32 fold degenerate

Corresponds to amino acid sequence:

trp lys lys gln lys asp lys

5' TGG AAA AAA CAG AAA GAC AAA

GGAGTG

Probe Sequence:

3' ACC TTT TTT GTC TTT CTG TT

C C T C A

Oligonucleotide Probe Pool LF2-5:

20mer 384 fold degenerate. Synthesized in four subpools of 96fold degeneracy.

Correspond to amino acid sequence:

gln gln ile tyr gly val val

5' CAG CAG ATC TAC GGN GTN GTN

AATT

Α

Probe Sequence:

3' GTC GTC TAG ATG CCN CAN CA

T T A A

T

LF2: 3' GTC GTC TAG ATG CCA CAN CA

T T A A

Т

LF3: 3' GTC GTC TAG ATG CCT CAN CA

TTAAA

Т

LF4: 3' GTC GTC TAG ATG CCG CAN CA

TTAAA

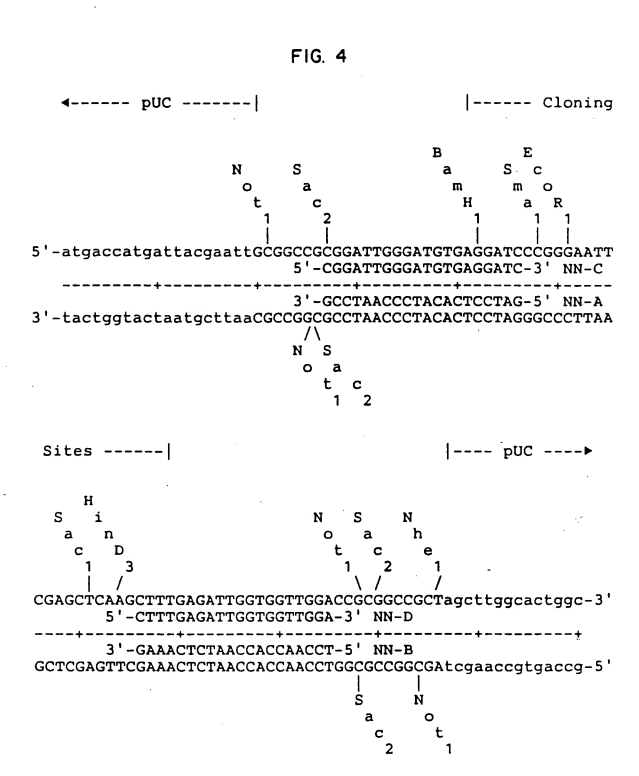
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FIG. 3

P24 AMINO ACID AND CDNA SEQUENCES

1	GCGGCCGCCGACGAGC	CATG(Met'	GTTGC ValAl	TGGG aGlv	AGCGACO Serasda	GCGGGGCGG AlaGlyArg	GCCCTG 50
	•	-28					,
51	GGGGTCCTCAGCGTGG	CTG	CCTGC	TGCA	ርፓርርጥጥባ	'GGTTTCA'	CACCTC 100
	GlyValLeuSerValVa	alCy	sLeuL	euHi	sCysPhe	GlyPheIl	eSerCy
101	TTTTTCCCAACAATAT	האדה:	ירים	СТСТ	· ATCCCAA		TOCAMO 1 E A
	sPheSerGlnGlnIle	rvrG	lvVal	Val Ti	rrGl vA c	10174611	TCCWIG 120
161	1 .	•			•		_
121	TACCAAGCAATGTGCCT	TTA	AAAGA	GGTC	CTATGGA	AAAAACAA	AAGGAT 200
	alProSerAsnValPro	•			•		
201	AAAGTTGCAGAACTGGA	CAAA	TTCTG.	AATTO	CAGAGCT	TTCTCATC	TTTTAA 250
	LysValAlaGluLeuGl	uAsı	nSerG	luPhe	ArgAla	PheSerSe	rPheLy
251		•		•	•	•	5Q
251	AAATAGGGTTTATTTAG	SACA	CTGTG	TCAGO	STAGCCT	'CACTATCT	ACAACT 300
•	sAsnArgValTyrLeuA	SpTi	nrVal	SerG]	lySerLe	uThrIleT	yrAsnL
201		•			•	•	•
301	TAACATCATCAGATGAA	GAT	GAGTA'	TGAAA	ATGGAAT	CGCCAAAT	ATTACT 350
	euThrSerSerAspGlu	Asp(GluTy	rGlub	ietGluS	erProAsn	IleThr
251		•		:		ŵ	•
221	GATACCATGAAGTTCTT	TCT	rtatg'	IGCTI	GAGTCT	CTTCCATC	TCCCAC 400
	AspThrMetLysPhePh	ieLei	TYTV	alLei	ıGluSer	LeuProSe	rProTh
401	ACTAACTTGTGCATTGA	CTA	TGGA	. הראי	ייירה א א הייי	CCAATGCA	TCATAC 4EA
	rLeuThrCysAlaLeuT	hrAs	nGlv	SerIl	eGluVa	lGlnCvsM	etileP
	•	•					
451	CAGAGCATTACAACAG	CAT	GAGG	ACTTA	TAATGT	ACTCATGG	GATTGT 500
	roGluHisTyrAsnSer	HisA	ArgGl	yLeuI	leMetT	yrSerTrp	AspCvs
	•	•		•		•	
501	CCTATGGAGCAATGTAA	LACGI	TAACT	CAACO	AGTATA	TATTTTAA	GATGGA 550
	ProMetGluGlnCysLy	sArg	gAsnS	erThr	Serlle	TyrPheLy	sMetGl
	• • • • • • • • • • • • • • • • • • • •	•		•		•	150
221	AAATGATCTTCCACAAA	(AAA)	racag'	TGTAC	CTCTTAG	CAATCCAT	TATTTA 600
	uAsnAspLeuProGlnI	ysIl	leGln	CysTh	rLeuSe	rAsnProL	euPheA
601			nmc			•	•
001	ATACAACATCATCAATC	ATT	TIGAC	AACCI	GTATCC	CAAGCAGC	GGTCAT 650
	snThrThrSerSerIle	irter	Leurn	rrnrc	Aerreh	roserser	GIAHIR
651	TCAAGACACAGATATGO	' እ ር ጥባ	P 3 TP 3 C		CCAMMA		
001	SerArgHisArgTyrAl	AL LA	TATAC	CATA	DEST	GCAGTAAT Alaualai	TACAAC 700
	- returnity this	anel	iiieb:	. 0116	rioLeu	wiangili	eThrTh 200
701	ATGTATTGTGCTGTATA	TGAZ	TGGT	• 4 ጥር ጥ	ጥርርጥጥ	TTAAAACA	
	rCysIleValLeuTyrM	letAs	nGl vi	MetTu	rAlaPh	e	MING /JU
751	TTTGAAAACTTGCATTC	STTTI	CCAA	AGGTO	AGAAAA	TAGTTTAA	GGATGA ANN
801	AAATAAAGTTTGAAATT	TTAC	SACAT'	TTGAA	AAAAA	AAAAAAA	AAAAA 850
851	AAAAAGCGGCCGC 86	3					



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FIG. 6 A

COMPARISON OF HT16 AND P24 cDNAs

```
V
               G
                 S D A G
             Α
                          R
                             Α
                              L G
HT16CGACGAGCCATGGTTGCTGGGAGCGACGCGGGGCGCGCCCTGGGGGTCCT 50
   P24 CGACGAGCCATGGTTGCTGGGAGCGACGCGGGGCGCCCTGGGGGTCCT 50
           V
             A G
                  S
         M
                    D A G
                          R
                            Α
                               L
                                 G V
      v
                   C
        V
                H
                     F
                         F
              L
                       G
                           I
                             S
                               C
                                  F
HT16 CAGCGTGGTCTGCCTGCACTGCTTTGGTTTCATCAGCTGTTTTTCCC 100
   P24 CAGCGTGGTCTGCCTGCACTGCTTTGGTTTCATCAGCTGTTTTTCCC 100
          CLLH
                  CFGF
                           I
                             S
                               C F
               v
                          T
         Y
                 Y
                   G
                     N
           G
HT16 AACAAATATATGGTGTTGTGTATGGGAATGTAACTTTCCATGTACCAAGC 150
   AACAAATATATGGTGTTGTGTATGGGAATGTAACTTTCCATGTACCAAGC 150
      IYGVVYGNVTFHVP
     V P L K E V L W
                     KKQKD
                                 K
                                   V
HT16 AATGTGCCTTTAAAAGAGGTCCTATGGAAAAAACAAAAGGATAAAGTTGC 200
   AATGTGCCTTTAAAAGAGGTCCTATGGAAAAAACAAAAGGATAAAGTTGC 200
   N V P L K E V L W K K Q K D K V A
         N S E
               F R
        E
                    Α
                      F
                        S S
                               K
HT16 AGAACTGGAAAATTCTGAATTCAGAGCTTTCTCATCTTTTAAAAATAGGG 250
   AGAACTGGAAAATTCTGAATTCAGAGCTTTCTCATCTTTTAAAAATAGGG 250
        ENSE
               F
                  RAF
                         S
                           S
                            FKNR
               S
                 G
                   S L
         D
           T V
                       T
                          Ι
                            Y
HT16 TTTATTTAGACACTGTGTCAGGTAGCCTCACTATCTACAACTTAACATCA 300
   TTTATTTAGACACTGTGTCAGGTAGCCTCACTATCTACAACTTAACATCA 300
      L D
           T
            V S G
                   S
                       T
                            Y
                     L
                          I
                                    S
         D
               E
                   E
                      S
                        P
                  M
                            I
                          N
HT16 TCAGATGAAGATGAGTATGAAATGGAATCGCCAAATATTACTGATACCAT 350
   P24
   TCAGATGAAGATGAGTATGAAATGGAATCGCCAAATATTACTGATACCAT 350
        D
           EYEM
                    E S P
                          N
                            Ι
          L
            Y
                   E
                     S
                       L
                         P
HT16 GAAGTTCTTTCTTTATGTGCTTGAGTCTCTTCCATCTCCCACACTAACTT 400
   GAAGTTCTTTCTTTATGTGCTTGAGTCTCTTCCATCTCCCACACTAACTT 400
              V L E
    K
     F
        F
            Y
                    SLP
                           S P
                          C
           N
             G
               S
                 I
                   E
                     V O
                            M
                              I
                                P
                                  E
HT16 GTGCATTGACTAATGGAAGCATTGAAGTCCAATGCATGATACCAGAGCAT 450
   GTGCATTGACTAATGGAAGCATTGAAGTCCAATGCATGATACCAGAGCAT 450
      LTNGSIEVQCM
                             IPE
```

FIG. 6B

	Y N S	H 1	R G I	LI	M Y	S W	D C	P M E	
HT16	TACAACA	GCCAT	CGAGGA	ATATT	ATGTA	CTCATO	GGATT	STCCTATGGA	500
					11111		11111	GTCCTATGGA	500
P24	TACAACA	GCCAT	CGAGGA	TTATA	ATGTA	CTCATO	GGATT	STCCTATGGA	500
	Y N S	H I	R G I	L I	M Y	S W	D C	P M E	300
	O C	KR	N S	T S	I Y	F	(M I	ENDT.	
HT16	GCAATGT.	AAACG!	TAACTC	ACCAG	TATAT.	ATTTTA	AGATG	SAAAATGATC	550
	111111				11111	!			330
P24	GCAATGT.	AAACG!	TAACTC	AACCAG	TATAT.	ATTTTA	LAGATG O	AAAATGATC	550
	Q C	K R	N S	T S	I Y	F F	ME	ENDL	330
	P Q	K I	Q C	T L	S	N P	L F	NTT	
HT16	TTCCACA	AAAAA!	TACAGTO	STACTC	TTAGC.	AATCCA	\ ጥጥ ል ጥጥ ነ	TAATACAACA	600
	111111	11111	11111		1111	11111			
P24	ŤŤĊĊĂĊĀ	AAAAA	TACAGTO	TACTO	TTAGC	AATCC	TTATT	TAATACAACA	600
	PO	K I	о с	TL	S	N P	L F	NTT	600
	S S I	I 1	. ີ ກັງ	: ב ב	I P	s s	GH	SRH	
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INTERNATIONAL SEARCH REPORT

International Application No PCT/US 89/03652

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ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

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